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(57) Abstract

A composite frozen confection product having ice confection material in contact with a layer of fat-based coating confection comprising a suspension of flavouring and sweetening solids in a fat component, characterised in that the fat component of the coating confection has: (a) a solids content index in each of the following ranges: 70-93% at -20°C; 65-93% at -10°C; 55-90% at 00C; 15-40% at 200C; 0-12% at 30°C; and 0% at 40°C; (b) a slip melting point in the range 23°C to 32°C; (c) a viscosity at 40°C 1/100 of at least 35 centipoise, and a viscosity at 40°C 1/2 CA of at least 30 centipoise; and in that the coating confection contains 30-70% by weight of the fat component and has a viscosity at 40°C of 1/2 at least 25 poise; 1/2 100 at least 2.0 poise and 7 CA at least 2.2 poise.

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COMPOSITE ICE CONFECTIONS AND PROCESSES FOR PREPARING THEM

The invention relates to composite ice confection products having layers or coatings (often called convertures) of fatty material, and to processes for preparing them.

Ice confections with fat-based coatings are wellestablished products. One known coating is chocolate, based on cocoa-butter. Other fats have been proposed for use in chocolate-like coatings, and some used. One fat that has been widely used is coconut oil.

Fat-based coatings can display properties that are
10 unsatisfactory to the consumer, the producer or both. Such
properties include the tendency to break, flake, or rub off
during production or consumption and to soften too much in
the hands.



Another important way in which fat-based coatings can be unsatisfactory is that they may give an unsuitable mouthfeel in the context of the product in which they are used. the one hand, such coatings may yield a lingering, cloying 5 mouthfeel, seeming to clog the teeth and oral surfaces, often also imparting bitterness to chocolate flavourings. other hand, coatings may fail to appear to contain appreciable chocolate flavouring or texture.

There are two main categories of chocolate coated 10 products relevant to this invention: enrobed products and Enrobed products are represented, for dipped products. example, by choc-ice bars (enrobed ice cream bars). cream bars are enrobed by passage through a curtain of molten coating material to cover the top and sides, and rollers are 15 used to coat the undersides during passage through the enrober. An airblower generally removes surplus coating material. Enrobing has been conventionally practised with cocoa butter based coatings, sometimes including vegetable fat blends.

Cocoa butter, a natural material which is expensive and 20 subject to unstable supply and market conditions, is also not ideal because it gives a coating which melts slowly onconsumption, and leaves a waxy impression when eaten as part The vegetable fat blends of a composite ice confection. have also suffered from comparable defects.

Dipped products are represented for example by many! chocolate coated confections on sticks, produced by dipping a core of material to be coated into a bath of molten Chocolate flavoured coatings for dipped coating material. products have been based on cocoa butter and on coconut oil. 30 Cocoa butter does not lend itself to satisfactory processing in this way: it is hard to obtain a thin uniform coated product. Coconut oil is in many ways a desirable material but, like cocoa butter, it produces an undesirably brittle coating, and again is subject to market and supply 35 difficulties.



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Another use of coatings based on fat compositions is in the coating of baked confectionery containers or wafers for ice confections. Fat coating of such baked articles is disclosed for example in GB 947,672 (Big Drum, Inc.). It has been found, however, that with the fat coatings in use there is a problem of loss of crispness in the baked product during storage for a reasonable time, due to water transfer from the adjacent ice confection.

According to the invention there are provided composite

10 frozen confection products having ice confection material in

contact with a layer of fat-based coating confection comprising

a suspension of flavouring and sweetening solids in a fat

component, where the fat component of the coating confection

has:

15 (a) solids content index (SCI) values (%) (as defined below) in each of the following ranges:

70-93 at -20°C; 65-93 at -10°C; 55-90 at 0°C; 15-40 at 20°C; 0-12 at 30°C; 0 at 40°C;

- (b) slip melting point in the range 230-320C; and
- (c) a viscosity at 40°C corresponding to measured values 25 (as defined below) as follows:

 $\eta_{100} \geqslant 35 \text{ centipoise;}$ $\eta_{CA} \geqslant 30 \text{ centipoise;}$

and wherein the coating confection contains for example 30-70% by weight of the fat component and has a viscosity at 40°C of 30 7 at least 25 poise; 7100 at least 2.0 poise and 7CA at least 2.2 poise. (In the foregoing and succeeding description reference is made to a number of parameters defined more closely by reference to test methods described hereinbelow.)

Preferably these fat compositions can also have SCI values (%) as follows,



at least 75%, e.g. 75-90 at -20° C; at least 75%, e.g. 75-85 at -10° C; at least 65%, e.g. 65-75 at 0° C.

They can also have an SCI of at least 57.5 at 10°C, e.g. 57.5-90%. Their preferred slip melting points are in the range 26° to 31°C.

We find that such fats give particularly good results in terms of manufacturing convenience and consumer acceptability owing to their lack of excessive brittleness. We find that they can be formulated into confectionery coating compositions of distinctly good flavour and mouthfeel on their consumption as part of a composite frozen confection. It can also be of advantage if the fat is one that gives a confectionery composition which in the test given below performs with a brittleness time of >45 seconds.

Normally such fats show a change in solids content over the last 5°C of their melting profile of at least 20% of their solids content at -20°C. Their SCI values at 10°C and 20°C can differ by, for example, at least about 25%, e.g. at least about 30%, and sometimes at least 35%. Their maximum rate of solids formation upon cooling is normally above 13-16% per minute, with the time taken to reach 50% solids normally less than 5-6 minutes. The maximum solids content achieved in \$20 minutes is normally in the range 70-90%.

Normally we prefer for ice confection use those fats which yield confectionery coatings of the following viscosities at 40°C when such coatings are formulated using 45% fat content by weight:

7₃ ≥ 25 poise, preferably 30-50 poise; 7₁₀₀ ≥ 2.0 poise, preferably 2.5-8.0 poise; 7_{CA} ≥ 2.2 poise, preferably 2.5-6.0 poise.

All these parameters are as defined by the test methods noted below or their equivalents.

Preferably these fats according to the invention are



made up by using interesterified mixtures of fats containing acyl groups derived from edible naturally-obtained oils or their equivalents. For example, we prefer to use interesterified blends of (a) lauric fats or oils and (b) non-lauric fats or oils in the weight proportions (a) up to 85%, e.g. 50%-75%; to (b) down to 15%, e.g. 45%-25% for preference: lower amounts of (a) of 15-50%, e.g. 25-40%, can also be used.

"Lauric fat or oil" means an edible oil or fat having
10 lauric and/or myristic acids as substantial fatty acyl
constituents, (e.g. constituting 40-50% or more of the acyl
groups) with the remaining fatty acyl content containing
very little unsaturation: examples are palm kernel oil
(preferred), babassu oil, coquito oil, coconut oil (also
15 preferred).

The term "non-lauric fat or oil" means an edible oil in which the principal fatty acyl content is not lauric acid, and for example in which there is a substantial amount of C₁₆ and/or C₁₈ unsaturated acyl groups, e.g. oleic, linoleic and/or linolenic acyl groups: the remaining bulk of the acyl groups is largely saturated, (e.g. C₁₆ and/or C₁₈ acids). Examples of such non-lauric oils are cottonseed oil, peanut oil, and particularly rapeseed oil (e.g. low-erucic rapeseed oil) and soyabean oil, and also palm oil.

When palm oil is used as the non-lauric oil its preferred proportions before interesterification are in the range 45%-25% by weight, especially 40%-35% by weight.

A particularly desirable fat in this category comprises a randomly interesterified blend of palm kernel oil (50%-30 65%) with palm oil (50%-35%), especially in the proportions 60:40.

Of course it is understood that in such interesterified blends other sources of acyl groups can be substituted for those quoted and for one another if their proportions and 35 types of acyl groups are sufficiently similar, without



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regard to their identical triglyceride combination or isomeric analysis. As an example, palm oil can in such compositions be substituted often by lard.

Another particularly preferred method of making up the fat according to the invention is to use an interesterified blend of palm kernel oil with rapeseed oil, in the proportions 15-60% to 85-40% respectively.

When randomly interesterified blends of a lauric fat or oil and a non-lauric oil other than palm oil, e.g. in proportions of 10-70% and 90-30% respectively, are used, the non-lauric oil or blend may be hardened (preferably selectively) to achieve the desired slip melting point.

(Mixtures containing palm oil are preferably not hardened because palm oil is already hard enough for most purposes.)

By selective hardening we mean exposure of the fat material to be treated to a catalyst which brings about isomerisation of the ethylenic bonds without substantial hydrogenation: considerable geometrical isomerisation (cis () trans) occurs. A suitable known catalyst for the purpose is a poisoned nickel catalyst (a fresh nickel catalyst gives much hydrogenation (unselective hardening) but little of the isomerisation required in selective hardening).

Preferably, because a desired slip melting-point in the product can be achieved more accurately, the interesterified fat is selectively hardened rather than the liquid oil or the blend. When a non-lauric oil other than palm oil is used, it can be one which is liquid or semi-liquid (e.g. of mushy consistency) at 20°C.

The fat-based coating confectionery compositions for use with ice confections comprise edible fat with a suspension therein of sweetening and (e.g. chocolate) flavouring agents in solid form, e.g. sugar and cocoa powder or substitutes therefor, and optionally also containing suspended milk solids.



It may be stated as a guide and without limitation that we prefer to use confectionery compositions containing from 30-70%, preferably 30-55% by weight fat, suitably at least 40%, e.g. about 45%. Emulsifiers, e.g. lecithin may be present as desired, e.g. in minor proportions about 0.2-1%, e.g. 0.5% up to 1%.

The coating confectionery can preferably have viscosities (at 40°C) as follows:

at least 25 poise, e.g. 30-45 poise;

¶ 100 at least 2.0 poise, e.g. 2.5-6.5 poise;

¶ CA at least 2.2 poise, e.g. 2.5-6.0 poise;

especially where they include milk solids. Especially where the coating confectionery comprises no milk solids, they can preferably have viscosities, (at 40°C) as follows:

One preferred example of a dark chocolate composition for use in composite frozen confections according to the invention, comprises 31% by weight of interesterified palm kernel oil and palm oil mixture (60:40 w/w), 24% cocoa mass, 39% sugar, with emulsifier and flavourant. A corresponding milk chocolate formulation can contain 33% by weight of a similar fat, and dried milk product in place of some of the sugar, to taste.

Such compositions among others can be used in accordance with the invention in processes for enrobing ice confections, or otherwise in any suitable way. It is certainly not excluded to use these compositions for dipping ice confections and so forming coatings otherwise than in an enrober, but it may then be necessary to apply blower air currents to thin the resulting coatings to a desired level. The compositions can also be used to coat wafer containers for subsequent filling with ice confections, e.g. cones.

35 Although the invention has been particularly illustrated so far by fats made up of edible oil blends as



described above, it is noted that very satisfactory fats and compositions can be made up by using blends or interesterified mixtures with other glyceride materials. For example, certain fats suitable for use according to 5 the invention are triglyceride mixtures comprising a major proportion of acyl groups derived from lauric fats or oils, or non-lauric fats or oils (optionally hydrogenated unselectively or else selectively hardened), and a minor proportion by number of acyl groups with substantially 10 short-chain lengths, e.g. up to C6. Preferably all or nearly all the short chain acyl groups are present in mixed triglycerides containing both short and long chains, e.g. So, for example, when palm kernel oil is used C>16 OF C>18. as a component of the fat, either alone, or in blend with, 15 or in interesterified mixture with another lauric fat or oil or a non-lauric fat or oil, the short chain acyl groups can be introduced by interesterification with for example glyceryl triacetate or glyceryl tributyrate or glyceryl trihexanoate, or they can be introduced by blending (and optional interesterification) with an acetylated long chain monoglyceride or diglyceride, or a long chain monoor di-glyceride acylated with short chain acyl groups.

Preferably the short chain acyl groups form up to about 20% by number of the total acyl groups of the tri25 glyceride mixture.

Where the short chain acyl groups are acetyl groups, they are preferably present at up to the equivalent of the number of acyl groups that would be provided by 5% by weight triacetin in tripalmitin or a mainly palmitic fat or oil, suitably up to 3% by weight, e.g. 2% by weight. Where the short chain acyl groups are butyryl or hexanoyl groups they are preferably present at up to the equivalent of the number of acyl groups that would be provided by 10% by weight glyceryl trihexanoate in tripalmitin or a mainly palmitic fat or oil, suitably up to 5% by weight.



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The use of hardened lauric oils gives fats with higher slip melting point than when untreated lauric oils are used. When hardened lauric oils are used, somewhat more short chains can be present than otherwise, since one effect of these is to lower the slip melting point.

The solids content index, brittleness time and viscosity referred to herein are measurements obtained as described below. The slip melting point is measured by standard procedures well known to those skilled in the filed to which this invention relates.

In this specification the solids content index (SCI) refers to a measurement obtained by the following or an equivalent test method: a sample of fat is placed in a low-temperature measurement cell of a wide-line nmr spectrometer calibrated for measurements of sample quantity, such as the "Quantity Analyser" commercially available from Newport Instruments, Newport Pagnell, Buckinghamshire, England. The sample is held at a required temperature for 10 minutes and then the percent solid fat is obtained from the instrument indication. This result is an SCI (%) referred to the test temperature.

The rate of crystallisation of a fat composition sample can conveniently be measured under conditions when the sample (at 32°C) is placed in an nmr cell of the same wideline nmr spectrometer at -20°C, and monitoring the signal level from the spectrometer at frequent intervals over about 20 minutes, intercalated with frequent temperature measurements using a temperature probe inserted whenever required into the sample cell, and removed to allow nmr measurement. The maximum rate of solids formation achieved is the steepest tangent to the curve of solids content versus time obtained from these measurements, and is expressed in % solids per minute.

In this specification "the brittleness time" refers to a measurement obtained by the following or an equivalent



test method in which what is estimated is the speed at which a confectionery coating composition (containing the fat to be tested) becomes brittle on being used to coat a frozen confectionery article. The fat composition to be tested is 5 incorporated at 59.2% of a confectionery composition cont containing 25.7% icing sugar, 12.9% cocoa powder (of 10-12% fat content), 1.7% skim milk powder and 0.5% lecithin. The total fat content of the composition should be 60.6%. Any conventional formulation process can be used: desirable the composition is rendered to a final particle size of 17-19 /um by micrometer screw gauge, with most particles of 14-16 jum as measured by a Sheen gauge: by Coulter Counter (Trade Mark) measurement 75% of particles are desirably & 22 /um. The confectionery composition prepared this way is melted at 36°C for 20 minutes in a gently stirred jacketed vessel, and used to coat frozen ice confection (ice cream) blocks on sticks in the following way. The blocks are desirably of 30g weight and 100% overrun, stored at -26°C. taken from the store and immersed 9 seconds in moulds surrounded by water at 30°C, so that their surface temperature 20 rises to -5°C and their interior temperature to -15°C: blocks are then immediately dipped into the molten coating composition (immersion for about ½ second), withdrawn, and small sections of the coated block repeatedly cut off with a 25 relatively blunt implement at closely-spaced time intervals by any suitable method until the coating can be heard to give an audible crack on cutting. The purpose is to simulate the conditions under which such compositions and coatings



are bitten during consumption, and a test consumer can be used if desired. The time elapsed between the coating immersion and the propensity of the coating to give an audible crack on cutting is taken as the "brittleness time" as referred to in this specification.

In this specification the viscosity measurements η_z , $\eta_{ ext{100}}$ and $\eta_{ ext{CA}}$ are measurements obtained by the following or equivalent test methods. Viscosity measurements can be made on fat compositions or on coating compositions made therefrom. 10 Where a coating composition is characterised, the viscosity measurement is made on the composition itself. Where a fat composition is in question, the viscosity measurements are either made on and referred to the fat composition itself or made on and referred to a coating composition comprising 15 45% of 59.2% by weight of the fat. Normally, coating compositions of 45% fat are used as standard where the coating compositions are intended for use in enrobing ice confections, and compositions of 59.2% fat where they are intended for use by dipping processes.

The parameters η_3 and η_{100} represent the apparent viscosity at the quoted temperature (e.g. 40°C) at shear rates of respectively 3 sec-1 and 100 sec-1. parameter $\mathcal{I}_{ exttt{CA}}$ represents the Casson plastic viscosity. $\eta_{
m 100}$ and $\eta_{
m CA}$ are measured, for example, using a Haake Rotovisko MVl (Trade Mark) viscometer with a rotating The fat or coating compositions to be inner cylinder. measured are desirably melted to a temperature 150 above the slip melting point and then equilibrated 15 minutes at the temperature of measurement. When measurements are 30 performed on couverture compositions these are pre-sheared 15 minutes at the measuring temperature at about 40 sec-1. For good order, apparent viscosity readings are taken through a range of machine shear rate settings in both ascending and descending senses, to determine the apparent viscosity 35 at the specified shear rate via determination of the shear stress for a given machine setting.



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The parameter $\eta_{\rm CA}$ (Casson plastic viscosity) is determined (e.g. using a similar viscometer to that mentioned above), but according to the method of N. Casson (Rheology of Disperse Systems, Pergamon Press, London 1959).

The parameter η_3 is measured for example using a Deer Rheometer PDR81 (Trade Mark) having an inner rotating cylinder or bob lowered onto a temperature-jacketed outer stationary cylinder containing the sample composition. The inner cylinder or bob is driven at a controlled shear stress and the shear rate induced is measured: the apparent viscosity at a 3 sec⁻¹ shear rate is determined from the results obtained at a succession of shear stresses.

The coating compositions described above have particularly desirable properties for use as enrobing convertures. That is, in coating compositions containing fat, cocoa powder or a substitute therefor, and sugar, optionally with a milk product and optionally with additives such as emulsifiers, these fats make compositions having desirable properties for use in the enrobing processes of ice confection coating, together with satisfactory organoleptic qualities. They are suitably formulated for example with viscosity parameter $\eta_{100} \ge 35$, preferably ≥ 50 , and parameter $\eta_{CA} \ge 30$, preferably ≥ 35 .

We find that certain of the preferred fats in this
category (e.g. 60:40 interesterified palm kernel oil:palm
oil) not only give superior processing characteristics in
enrobing processes, with reduced brittleness, but the
resulting coatings have even better mouthfeel than that of
coatings based on cocoa butter (which leads to a very
brittle result).

The invention is now described for illustration but without limitative effect by means of the following Examples in conjunction with the foregoing description.



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Example 1

Blends of 60% PKO with 40% palm oil were made from neutralised, fully refined dried oils. 2-2.5 kg batches of the blends were interesterified in a batch process, in a flask fitted with a vigorous stirrer, nitrogen inlet, and temperature control probe. The blends were heated to 110°C under N₂, stirred vigorously and sodium methoxide catalyst (0.2% on the oil) added into the vortex. Stirring was maintained for 30 minutes.

The product was washed with 1% acetic acid solution, followed by water, to remove catalyst and soap, and thoroughly dried. Completion of interesterification was checked by triglyceride GLC analysis.

The dried, interesterified oils were bleached with 2% 15 bleaching earth at 110°C under nitrogen for 30 minutes. 1% kieselguhr was added to the cooled blends, which were filtered.

650g batches of the interesterified blends were deodorised for 4 hours at 180°C under vacuum, and 0.1% of lecithin was added to each deodorised batch, as antioxidant.

Batches of this interesterified blend had slip melting points of 29.5-31°C and had solids content melting profiles and crystallization curves as shown in Figures 1 and 2, when analysed under the standard test conditions described above.

The maximum rate (%/minute) of solids formation under the standard test conditions in the batches was 16-23%/ minute, with 4-5 minutes required on cooling to reach 50% solid. They had periodate values less than 4, usually 30 0.5 or less, iodine values of about 31-32, and hydroxyl values about 8-10. The refined oils used in their preparation had FFA contents less than about 0.5 and soap values less than about 0.01.

The percentages solids present at various temperatures 35 were as follows:



-20°C	79-85%;
-10°C	77-82%;
Ooc	67-72%;
10°C	48-58%;
20°C	16-36%;
25 ⁰ C	5-20%;
30°C	0-5%;
35 ⁰ C	0%.

Excellent enrobedice creambars are prepared by per se

10 conventional handling techniques using these products. Processing
was easier (shorter setting-up time and less brittle) than
with a conventional converture based on cocoa-butter.

Examples 2 and 3

(A) Preparation of fat compositions

O.8N NaOH. After a water wash, the oil was boiled with 1.5% its weight of 6N soda ash and 1% by weight of a sodium silicate solution for 30 minutes at 105°C, then washed free of soap and dried. Bleaching was carried out 20 with 1% Tonsil Standard FF (Trade Mark) bleaching compound 20 minutes at 105°C under vacuum. The neutralised bleached oil was doedorised 30 minutes at 180°C.

The rapeseed oil so treated was blended with fullyrefined palm kernel oil in the weight ratio 40 rapeseed oi
to 60 palm kernel oil in a first case (Example 2) and 50
to 50 in a second case (Example 3). Batches of the blends
were interesterified continuously with 0.05% by weight
sodium catalyst at 125°C for 4 minutes. The product oil
was then added to a vessel partly filled with diluted
phosphoric acid (to neutralise excess sodium). The
treated product oil (interesterified blends) were washed
with 0.2N NaOH and water; when free of soap the blends
were dried and filtered over 0.2% Hyflow (Trade Mark) filter
aid.



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Batches of the blends were selectively hardened under agitation in an autoclave fitted with six-blade turbine stirrer and four baffles. Hydrogen was blown into the headspace of the autoclave. Hardening was carried out at 180°C with 0.5% by weight of sulphur-poisoned nickel catalyst at 3-5 kg/cm² H₂ pressure. The blends were hardened to slip melting-points of 26.8°C (Example 2) and 26.7°C (Example 3). The selectively hardened interesterified blends were neutralised, freed of soap, dried, bleached and doedorised in similar manner as before. At 60°C, 0.1% fresh lecithin was added to each as antioxidant.

The properties of the fats prepared as described above were found to conform desirably with those described above.

15 (B) Preparation of confectionery coating compositions and ice confections

Using normal refining and conching procedures, the fat compositions prepared according to the invention in Examples 2(A) and 3(A) were made up into confectionery coating compositions containing 43.6% of the respective fat, 39.9% sugar, 16% cocoa powder (10/12% fat) and 0.5% lecithin, with optional flavourant to taste. These compositions were found to have highly desirable physical properties when used to coat ice confections (e.g. using an ice-briquette enrober and 40°C coating temperature) and the flavour and mouthfeel characteristics of the products were found to compare satisfactorily with those of products based on 20% cocoa-butter fat.

Example 4

(i) Further interesterified blends of palm kernel oil (PKO) and palm oil (PO) according to the invention were prepared as follows.

A crude blend (60 PKO:40 PO) was made up in a vacuum bleacher, vacuum dried, neutralised with 6N soda ash,



washed and dried. The dried product was bleached at 95-100°C for 30 minutes with 1% C300 bleaching earth, filtered and transferred to a clean vessel. There followed interesterification with 0.3-0.4% sodium methoxide catalyst, and the treated blend was washed, vacuum dried, and post-refined with 1% AA bleaching earth at 95-100°C for 30 minutes. After filtration and deodorisation 0.1% lecithin was added.

(ii) The interesterified blends produced in Example 4(i)
10 were used to make up confectionery coating compositions
according to the invention, which were used to enrobe ice
confection briquettes as described in Examples 2 and 3.

The compositions (Examples 4(A), 4(B), 4(C), and 4(D), were of the following constituents:

15		<u>4A</u>	<u>4B</u>	<u>40</u>	<u>4D</u>
	Cocoa solids (non-fat)	14.2%	16.3%	5.1%	5.1%
	Milk solids (non-fat)	-	-	18.8%	18.8%
	Sugar	39.9%	39.0%	30.6% ·	30.6%
	Lecithin	0.5%	0.5%	0.5%	0.5%
20	Cocoa butter	1.8%	13.2%	0.7%	6.5%
	Butterfat	_ .	-	3.5%	3-9%
	Interesterified fat blend of Example 4(i)	43.6%	31.0%	40.8%	34.6%
	(Total fat content)	(45.4%)	(44.2%)	(45.0%)	(45.0%)

All showed excellent processing and consumer-appreciable properties. In comparative evaluations, the interesterified blends according to embodiments of this invention were shown to give superior results to otherwise equivalent blends of PKO and PO without interesterification, (such comparative blends lack the physical properties of the fats of this invention specified above). Compositions 4B and 4D had particularly desirable processing and consumer-appreciable characteristics.



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The invention is not limited to the use of the materials particularly mentioned above. Further particular examples of fat compositions falling within the general classes mentioned above, which are useful, are interesterified blends of palm kernel oil and cottonseed oil (15%:85%), selectively hardened to slip melting point 25°-29°C; of palm kernel oil and low-erucic rapeseed oil (15%:85%), selectively hardened to slip melting point 26°C; and of palm kernel oil and soyabean oil (15%:85%), selectively hardened to slip melting point 25°-29°C. In each case as in the other cases described above the non-lauric oil can in part or even wholly be substituted by alternative materials, e.g. other non-lauric oils, or tallow olein, or palm olein.



CLAIMS:

1. A composite frozen confection product having ice confection material in contact with a layer of fat-based coating confection comprising a suspension of flavouring and sweetening solids in a fat component, characterised in that the fat component of the coating confection has:

(a) a solids content index in each of the following ranges:

70-93% at -20°C; 65-93% at -10°C; 55-90% at 0°C; 15-40% at 20°C; 0-12% at 30°C; and 0% at 40°C;

- (b) a slip melting point in the range 23°C to 32°C;
- (c) a viscosity at 40° C η_{100} of at least 35 centipoise, and a viscosity at 40° C η_{CA} of at least 30 centipoise; and in that the coating confection contains 30-70% by weight of the fat component and has a viscosity at 40° C of η_{3} at least 25 poise; η_{100} at least 2.0 poise and η_{CA} at least 2.2 poise.
- 2. A composite frozen confection product according to Claim 1, characterised in that the fat component has solids content index of

at least 75% at -20° C; at least 75% at -10° C; and at least 65% at 0° C.

- 3. A composite frozen confection product according to Claim 1 or 2, characterised in that the fact component has a solids content index in the range 57.5-90% at 10°C.
- 4. A composite frozen confection product according to Claim 1,2 or 3, characterised in that the solids content indices of the fat component at 10°C and at 20°C differ by at least about 25%.



- 5. A composite frozen confection product according to Claim 1,2 or 3, characterised in that the solids content indices of the fat component at 10°C and at 20°C differ by at least about 30%.
- 6. A composite frozen confection product according to Claim 1, characterised in that the fat component has solids content index in each of the following ranges:

at -20°C, about 79-85%, at -10°C, about 77-82%, at 0°C, about 67-72%, at 10°C, about 48-58%, at 20°C, about 16-36%, at 25°C, about 5-20%, at 30°C, about 0-5%, at 35°C, 0%.

- 7. A composite frozen confection product according to any of claims 1-6, characterised in that the coating confection has viscosities at 40°C in the ranges 7 3 30 to 50 poise; 100 2.5 to 8.0 poise; 7 CA 2.5 to 6.0 poise.
- 8. A composite frozen confection product according to Claim 1-7, characterised in that the coating confection contains at least 40% w/w of the fat component, for example up to 55% w/w.
- 9. A composite frozen confection product according to Claim 1-8, characterised in that the fat component comprises an interesterification product of a fat mixture containing a lauric fat or oil.
- 10. A composite frozen confection product according to Claim 9, characterised in that the fat component comprises an interesterification product of a fat mixture containing a lauric fat or oil and a non-lauric oil.



- 11. A composite frozen confection product according to Claim 9, characterised in that the fat component comprises an interesterification product of a fat mixture containing a lauric fat or oil and a glyceride containing short-chain $(C_{\leq 6})$ acyl groups.
- 12. A composite frozen confection product according to Claim 9, characterised in that the fat component comprises an interesterification product of a fat mixture containing a lauric fat or oil, and glycerides containing long chain $(C_{\geq 16})$ acyl groups and short chain $(C_{\leq 6})$ acyl groups.
- 13. A composite frozen confection product according to Claim 9, characterised in that the fat component comprises an interesterification product of a fat mixture comprising 25-85% lauric fat or oil.
- 14. A composite frozen confection product according to Claim 9, characterised in that the fat component comprises an interesterification product of palm kernel oil (50-65%) and palm oil (50-35%) or low-erucic rapeseed oil (50-35%).
- 15. A composite frozen confection product having ice confection material in contact with a layer of fat-based coating confection, wherein the fat component of the coating confection comprises an interesterification product of pakernel oil (60-65%) and palm oil (40-35%).
- 16. A composite frozen confection product having ice confection material in contact with a layer of fat-based coating confection, wherein the fat component of the coating confection comprises an interesterification product of palm kernel oil or coconut oil (15-85%) with low-erucic rapeseed oil, cottonseed oil, soyabean oil or groundnut oil (85-15%).



- 17. A process for producing a composite frozen confection product which comprises contacting together ice confection material and a layer of a fat-based coating confection comprising a suspension of flavouring and sweetening solids in a fat component, characterised in that the ice confection is contacted with a fat-based coating confection comprising a fat component having:
- (a) a solids content index in each of the following ranges:

70-93% at -20°C; 65-93% at -10°C; 55-90% at 0°C; 15-40% at 20°C; 0-12% at 30°C; and 0% at 40°C;

- (b) a slip melting point in the range 23°C to 32°C;
- (c) a viscosity at 40°C η_{100} of at least 35 centipoise, and a viscosity at 40°C η_{CA} of at least 30 centipoise; and containing 30-70% by weight of the fat component and has a viscosity at 40°C of η_3 at least 25 poise; η_{100} at least 2.0 poise and η_{CA} at least 2.2 poise.
- 18. A process according to Claim 17, characterised in that said ice confection material is coated with said fat-based coating confection at a temperature of about 40°C.
- 19. A process according to claim 17, characterised in that confection material is contacted with a baked container having a coating of said fat-based coating confection.
- 20. A process according to Claim 17,18 or 19, characterised in that said fat-based coating confection is as defined in any one of claims 2 to 16.



INTERNATIONAL SEARCH REPORT International Application No PCT/GB 80/00036 1. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) \$ According to International Patent Classification (IPC) or to both National Classification and IPC Int. Cl.³: A 23 G9/02; A 23 D 5/00 II. FIELDS SEARCHED Minimum Documentation Searched 4 Classification System Classification Symbols Int.C1.3 A 23 G 9/02; A 23 G 1/00; A 23 G 3/00; A 23 G 9/24; A 23 G 9/28; A 23 G 9/00; A 23 D 5/00; A 23 D 3/00 Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched 5 III. DOCUMENTS CONSIDERED TO BE RELEVANT 14 Citation of Document, 16 with indication, where appropriate, of the relevant passages 17 Category • Relevant to Claim No. 18 US, A, 3333968, published August 1st, 1967 1,8-10,12, see column 1; column 2, lines 13-65, examples 1-4, column 1, lines 54-58, 16-18, 20 column 3, lines 39-46, R.S. Bell et al. FR, A, 1459649, published November 18, 1966 see page 4, column 1, alineas 3,4, page 1-4,6,8-10,12, 13, 17, 5, column 1, alineas 3,4, tables 1-5, 18,20 abstract, National Biscuit Comp. FR, A, 2121601, published August 25, 1972 1,8-10,12, see page 1, lines 4-6, columns 1,3-7, 13,15-17, 10,11,15, page 4, line 30, examples 4, 19,20 5,9-14, Unilever BE, A, 623770, published February 15, 1963 15-17, 19, 20 see page 9, alineas 2 and 3, Big-Drum FR, A, 1414086, published by Bulletin 1, 11, 12 Officiel de la Propriété Industrielle No. 42, 1965 see examples 1-4. abstract. The Procter & Gamble GB, A, 1017480, published January 19, 1966 Special categories of cited documents: 18 A" document defining the general state of the art "E" earlier document but published on or after the international filing date "P" document published prior to the international filing date but on or after the priority date claimed "L" document cited for special reason other than those referred "T" later document published on or after the international filing date or priority date and not in conflict with the application, but cited to understand the principle or theory underlying the invention to in the other categories "O" document referring to an oral disclosure, use, exhibition or "X" document of particular relevance IV. CERTIFICATION Date of the Actual Completion of the International Search 3 Date of Mailing of this International Search Report : June 25, 1980

July 7, 1980

International Searching Authority & European Patent Office

Signature of Authorized Officer 20

G.L.M. KRUYDENBÈRO

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BNSDOCID: <WO____ 8001749A1 I >

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	SERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 10
This inter	national search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons: im numbers because they relate to subject matter 12 not required to be searched by this Authority, namely:
2 Clai	im numbers, because they relate to parts of the international application that do not comply with the prescribed require- nts to such an extent that no meaningful international search can be carried out ¹³ , specifically:
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VIJ 0	BSERVATIONS WHERE UNITY OF INVENTION IS LACKING 11
	ernational Searching Authority found multiple inventions in this international application as follows:
of t	all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims the international application.
2 As	s only some of the required additional search fees were timely paid by the applicant, this international search report covers only ose claims of the international application for which fees were paid, specifically claims:
3. No	o required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to e invention first mentioned in the claims; it is covered by claim numbers:
	on Protest he additional search fees were accompanied by applicant's protest. o protest accompanied the payment of additional search fees.

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V. 1 OB:	SEDVATIONS WHERE CERTAIN OF ANY MANY
	SERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 10
1. Clair	national search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons: n numbers, because they relate to subject matter 12 not required to be searched by this Authority, namely:
2. Clain ment	n numbers, because they relate to parts of the international application that do not comply with the prescribed require- s to such an extent that no meaningful international search can be carried out ¹³ , specifically:
VI. OBS	SERVATIONS WHERE UNITY OF INVENTION IS LACKING 11
This Intern	ational Searching Authority found multiple inventions in this international application as follows:
1. As all of the	required additional search fees were timely paid by the applicant, this international search report covers all searchable claims international application.
2. As on those	nly some of the required additional search fees were timely paid by the applicant, this international search report covers only claims of the international application for which fees were paid, specifically claims:
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J. No rec	quired additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to vention first mentioned in the claims; it is covered by claim numbers:
Remark on I	Protest
	dditional search fees were accompanied by applicant's protest.
No pre	otest accompanied the payment of additional search fees.

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